# EUROPEAN PATENT APPLICATION published in accordance with Art. 158(3) EPC

(43) Date of publication: 10.05.2006 Bulletin 2006/19

(21) Application number: 03776031.1

(22) Date of filing: 02.12.2003

(51) Int Cl.: H01L 21/304 (1974.07)

(86) International application number: PCT/JP2003/015431

(87) International publication number: WG 2005/015625 (17.02.2005 Gazette 2005/07)

(84) Designated Contracting States:

AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
HU IE IT LI LU MC NL PT SE SI SK TR

(30) Priority: 12.08.2003 JP 2003292523

(71) Applicant: S.E.S. Company Limited Ome-shi, Tokyo 198-0023 (JP)

(72) Inventors:

 NAKATSUKASA, Katsuyoshi, S.E.S. CO., LTD.
 Asakuchi-gun,

Okayama 719-0302 (JP)

OGASAWARA, Kazuhisa.

S.E.S. CO., LTD. Asakuchi-gun, Okayama 719-0302 (JP)  SAKAIHARIA, Yoshiaki, S.E.S. CO., LTD. Asakuchi-gun, Okayama 719-0302 (JP)

 HARUKI, Yoshihiro, S.E.S. CO., LTD.
 Asakuchi-gun.

Okayama 719-0302 (JP)

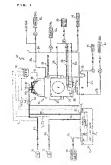
KAWATE, Munenori.
 S.E.S. CO., LTD.
 Asakuchi-gun.

Okayama 719-0302 (JP)

(74) Representative: HOFFMANN EITLE Patent- und Rechtsanwälte Arabeliastrasse 4 81925 München (DE)

### (54) METHOD OF PROCESSING SUBSTRATE AND SUBSTRATE PROCESSING APPARATUS

A substrate processing apparatus 10 including a vapor generating unit 37, which generates a mixed gas consisting of an organic solvent vapor and an inert gas by bubbling the inert gas in the organic solvent; support means for supporting a plurality of substrates to be vertically arranged in parallel at equal pitches; a processing vessel 15 which accommodates multiple substrates supported by the support means: a lid 30 for covering the upper opening of the processing vessel; jet nozzles 33 provided in the lid 30; and first piping 3712, 342, 3421, and 34,2 which causes the vapor generating unit and the jet nozzles to communicate with each other. In the substrate processing apparatus 10, the first piping and the et nozzles are respectively equipped with heaters, and the heaters are controlled by means of dry gas containing organic solvent mists of submicron size being emitted from the jet nozzles. According to the invention, Since micro-size organic solvent vapor is used, the substrate processing method and apparatus of the invention ensures not only high-quality surface processing but also the reduction of processing time.



### Description

#### TECHNICAL FIELD

[0001] The present invention relates to a substrate processing method and a substrate processing apparatus designed to perform drying processing on substrate surfaces such as a semiconductor water, a liquid crystal display device substrate, a recordind disk substrate and a mask substrate most efficiently and with a hind derive of outsite.

#### BACKGROUND ART

10

[0002] In the process of cleaning the surface of various substrates, for example, a semiconductor water (hereinatter referred to as water), is cleaned by applying chemicals to the surface thereof, and then rineed with a processing solution such as pure water, and thereafter cried with an organic solvent such as isopropyl alcohol (hereinatter referred to as IPA). More specifically, the processing includes a process, in which, after it has been cleaned with chemicals and pure water, the water is exposed to IPA vapor to condense IPA on the wafer surface to substitute the IPA alchering to the water with pure water, and certain contaminants such as particles are washed out from the wafer surface, and a drying process by which IPA is vaporized to dry the wafer surface, in the drying process, even the smallest water droplet remaining on the wafer surface can form a water mark on the wafer surface, thereby adversely affecting the quality of the wafer. Therefore, in the semiconductor production process, it is necessary to ensure that contaminants do not adhere to the wafer. Various methods and apparatuses for processing substrate surfaces of wafers and the IIKe, including measures dealing with contaminants, have been devised and put to practical use, and such method and apparatus for processing the substrate have been disclosed in several patent documents, including Japanese Laid-Open Patent No. 2001;27:188 (see Fig. 1 and left column of page 5 to left column of page 6) and Japanese Laid-Open Patent No.

6 [0003] Hereafter, the substrate processing apparatus disclosed in Japanese Laid-Open Petent No. 2001:271188 will be described with reference to Figs. 10 and 11. Fig. 10 is a sectional view of the substrate processing apparatus described in Japanese Laid-Open Patient No. 2001:271188. A substrate processing apparatus 1 compreting a processing vessel 2, a processing solution introducing pipe 3, a vapor generation unit 4, a processing solution drain unit 5, and heated-solvent supply devices 6 and 6. The processing vessel 2 houses the substrates (for example, water) to be processed. The processing solution introducing pipe 3 supplies the processing solution for example, pure water) into the processing solution.

The processing solution introducing pipe 3 supplies the processing solution (for example, pure water) into the processing vessel 2. The vapor generation unit 4 houses the organic solvent solution (for example, IPA). The processing solution drain unit 5 drains the processing solution from the processing vessel 2. The heated-solvent supply devices 5 and 6' supply the heated organic solvent into the vapor generation unit 4. The plant substrates are conveyed into the processing vessel 2 while vertically standing in parallel af equal intervals, and substrate surface processing is performed.

[0004] In the substrate processing apparatus 1, the surface processing of various substrates, e.g., a semiconductor wafer W (hereinafter referred to as wafer W) is performed in accordance with the following processes.

# 1) Wafer Conveying Process

Q [0005] A lid 2<sub>1</sub> of the processing vessel 2 for keeping pure water J in a standby state in the processing exparatus 1 is opened, and the pluris wafers W are conveyed to an inner vessel 2<sub>2</sub> of the processing vessel 2, and thrown and dipped into the pure water J, and the lid 2<sub>1</sub> is closed. Then, an inert gas such as a nitrogen gas is supplied from an inert gas supply pipe 8<sub>1</sub> into the processing vessel 2, to replace air in the processing vessel 2.

#### 45 2) Drying Process

[006] Then, after the water W is washed or rinsed, bubbling nitrogen gas N<sub>e</sub> is supplied to the vapor generation unit 4 to generate vapor of an organic solvent solution, e.g., IPA, and the vapor generated is conveyed from the discharge port 4, to the processing vessel 2, and the space above the pure water J therein is filled with the vapor,

[0007] Then, the on-off valve of an inner vessel drail pipe 5, is opened to drain the pure water J in the inner vessel 2,5 bit by bit through a flow control valve, and the liquid level of the pure water J gradually decreases to expose the water W from the upper end thereof to the liquid surface of the pure water J.

[0008] When the surface of water W is exposed white hying above the figuid surface of the pure water d, the IPA vapor in the processing vessel 2 comes into contact with the surface of water W. At this point, because the pure water J of the processing vessel 2 is substantially set at room temperature, the temperature of the water W nearly reaches room temperature also. Therefore, by coming into contact with the water W, the IPA vapor is rapidly cooled and condenses on the surface of the water W on the liquid surface. The condensed IPA reduces surface tension of the pure water that has addread to the water W. After the pure water later J is completely drained, linef das acts to replace the pure water that has addread to the water W. After the pure water later J is completely drained, linef das

is supplied to the processing vessel 2 from the inter gas supply pipe 8, in order to dry the surface of the water W by causing the IPA to evaporate.

3) Wafer Taking Out Process

[0009] Then, after the pure water J is drained from the inner vessel  $2_2$ , the substrate processing is completed by taking out the water W from the processing vessel 2 upon opening the lid  $2_2$ .

[0010] In the substrate processing apparatus 1, since the series of processes is performed in one closed processing viscel, the water never comes into contact with the atmosphere, and the adhesion of contaminants such as foreign particles and water marks as well to the water surface can be avoided white ensuring efficient processing thereof.

[0011] However, in recent years, it has become necessary to insert into the processing vessel wafers processed by this kind of substrate processing apparatus in large numbers as may be possible in order to increase processing efficiency. In some cases, the waters are simultaneously processed in the processing vessel in lot units ranging from 50 to 100 wafers, so that the space between wafers tends to become smaller, in addition, the diameter of wafers has increased from 200 mm to 300mm. Accordingly, while the generation of water marks can be suppressed in wafers of relatively small diameter, such as 200 mm or less, such is not the case for wafers having a diameter of as large as 300 mm. The processing efficiency of the conventional apparatus is therefore limited.

[0012] By examining water marks formed from verious angles, the inventors discovered that since the IPA vapor in dry gas is obtained by publicing the inert gas in IPA. I contains a large amount of liquid particles of minor IPA (pare interfer referred to as "mist") other than IPA gas with lower saturation concentration, and the size and mass of the mist is larger than those of histogen gas. Therefore, the IPA mist hardly passes through the narrow gaps between walers. Accordingly, when the diameter of the water becomes 300 mm, IPA substitution is not sufficiently performed because the IPA mist is hardly supplied to the water surface which is far from a supply port of the IPA mist. In other wards, in cases in the interfer supplied amount of IPA contained in the dry gas, the number of mists decreases as the size of the IPA mist increases. Conversely, the number of mists increases as the size of the IPA mist decreases. Further, when the IPA mist is large in size, the mase is heavy, thereby diminishing its moving speed. Therefore, in the drying process described in section (2) above, even if dry gas is supplied between the plural waters in the processing vessel, there is an imbalance between the number of water origidate. For example, when the number of IPA mist particles is smaller than the number of water droplets of the rinsing solution achiering to the water surface and the number of IPA is not substituted for some of the water droplets with the rinsing solution achiering solution achiering is the generation of water marks.

guostitude for some of the water dropues which interior termain, resulting in the generation of water marks. (D013) In addition, since it is enormous in size and heavy, the IPA mist hardly passes through the nerrow gaps between waters. Therefore, in wafers having a diameter as large as 300 mm, almost all the IPA mists address to the water surface with its relatively father from the supply port of the IPA mist before the IPA mist serve the water surface which is relatively father from the supply port. Accordingly, in the wafer surface distantly located from the supply port, the number of IPA mists supplied to the wafer surface ener the supply ont of the IPA mist is more than required while the number of IPA mists supplied to the wafer surface distantly located from the supply port is sufficient, resulting in the situation where IPA is not sufficiently established for the missing solution adhering to that part of the wafer surface which is dietantly located from the IFA mist supply port, therefore the surface which is dietantly located from the IFA mist supply port, thereby resulting in the generation of water marks.

[0014] The state in which IPA is substituted for water drojets will be described with reference to Fig. 11. Fig. 11 is a sectional view exhematically showing the relationship between the IPA mist and water drojets of the rinning solution (hereinafter referred to as DIW) adhering onto the wafer W in the drying process. In the drying process described in section (2) above, as shown in Fig. 11 (a), a mixture of a nitrogen gas Ng and the IPA vapor containing large-size IPA and stil (tiquid) is supplied into the processing vessel, for application to the spaces between wafers W. As shown in Fig. 11 (b), although the IPA vapor is intended to replace DIW, the moving speed is slow due to the large size of the IPA mist, and because many wafers (50 to 100 wafers) with a diamater of as large as 300 mm are similarinaeously processing in the processing vessel, the number of IPA mist is likewise limited, and thus there are occasions when the IPA mist does not reach all the DIWs. Thus, the IPA mist advisers to that portion of the wafer surface has the protion of the wafer surface which is terther from the dry gas supply port. Accordingly, as shown in Fig. 11(c), DIW remains on the wafer surface, which is resulted or water marks.

[0015] In the substrate processing apparatus disclosed in Japanese Laid-Open Patent No. H11-191569, the organic solvent is healed and evaporated to generate the mixture of organic solvent rough and her days in the evaporation vessels without bubbling the inert gas in the organic solvent, and then the mixed gas is heated and kept constant while it is diluted with another inert gas originating from a piping, and the mixed gas is then emitted through a jet nozzle. In the above-mentioned substrate processing apparatus, the organic solvent vapor originating and emitted from the piping and the nozzle completely becomes the mixed gas to be used. In such event, because the size of the gaseous organic solvent molecule is much smaller than that of the mixt, the problem related to the generation of water marks out or

use of the organic solvent mist does not arise.

[0016] However, even if the substrate processing is performed with the organic solvent vapor which completely pecentes the mixed gas, because the concentration of organic solvent uppor in the dy gas does not go beyond the saturation point, the absolute amount of organic solvent in the dry gas is rather small. Accordingly, the spread of the organic solvent vapor to every corner of the large substrate to replace the moisture of the substrate surface takes considerable time and therefore the substrate processing appearants disclosed in Japanese Laid-Open Patient No. H11-191549 does not address the need to decrease and substantially eliminate the number of water marks formed on the substrate surface while increasing the speed of carrying out the drying processing.

[0017] Although the term "vapor" generally means "gas" in the technical field of substrate processing, "vapor" can also refer to a gas containing "micro liquid particles (mistr)", such as the dry gas previously referred to. Accordingly, both meanings shall apply when "vapor" is mentioned in the present description and the claims.

### DISCLOSURE OF THE INVENTION

- 15 [0018] As a result of various studies conducted by the inventors considering the abovementioned prior references, the inventors discovered that when the size of the mist constituting the organic solvent vapor becomes extremely smaller, the number of organic solvent mist particles can be increased without the need of increasing the amount of organic solvent to be consumed. Further, eithough the surface area of a particular mist diminishes, the whole surface area expressed by a summation of the mists can be increased by causing the number of mists to increase. When the organic solvent vapor in which the surface area of the entire number of mists is increased is injected to the substrate surface, the organic solvent vapor can cover all water droplets adhering to the substrate, so that the organic solvent can efficiently be substituted for the water droplets.
- [0019] The first object of the Invention is to provide a substrate processing method in which not only high-quality substrate surface processing is realized but in which processing time is also shortened by using dry gas containing micro-size organic solvent mist.
  - [0020] The second object of the invention is to provide a substrate processing method in which not only high-quality substrate surface processing is realized but in which the processing time is shortened by adjusting the concentration of micro-size organic solvent mist in the dry gas.
- [0021] The third object of the invention is to provide a substrate processing apparatus in which not only high-quality substrate surface processing is realized but in which the processing time is shortened by easily forming the dry gas containing micro-size corpanies cohert micro.
  - [0022] The fourth object of the invention is to provide a substrate processing apparatus in which not only high-quality substrate surface processing is realized but in which the processing time is shortened by adjusting the concentration of micro-size organic solvent mist in the dry age.
- 50023] In crider to stain the above objects, the substrate processing method according to claim 1 of the invention refers to a substrate processing method in which the surface of a substrate is dried by injecting it with dry gas containing a mixture of an organic solvent vapor and an inert gas, characterized in that the organic solvent vapor contains mists having submicron size.
  100241. According to the substrate processing method described in claim 1, since the size of the mist contained in the
- 90 organic solvent vapor is reduced to submicron size, the number of perticles of the organic solvent mist can be increased without increasing the amount of organic solvent to be consumed. Further, elthough the surface area of the individual mist diminishes, the whole surface area which constitutes a summation of the surface arease of the individual mists can be increased by causing the number of mists to increase. Therefore, since a large amount of mists of submicron size can be injected onto the substrate surface, a large amount of organic solvent mist of submicron size is efficiently substituted for the rinsing solution adhering to the substrate. As a result, even if multiple substrates with a large diameter are loaded in the processing vessel, the processing time can be stortened since the mists of submicron size can rapidly cover the gaps between substrates, while the rate of efficiency of the drying processing improves, and the generation of water marks in the substrate surface can be largely avoided or substantially eliminated. Further, since adhesion of particles is eliminated and the speed of drying processing is increased, re-adhesion of particles or be prevented.
- (0025) The substrate processing method according to claim 1 of the invention is characterized in that the dry gas is a mixture of finer gas and organic solvent vapor formed by bubbling the linert gas in the organic solvent in a vapor generating unit, wherein the temperature of the vapor generating unit is set at T<sub>1</sub>, the temperature of the mixed gas containing the inert gas and the organic solvent is set at T<sub>2</sub> from the vapor generating unit to a jet nozzle, and the temperature of the dry gas emitted from the jet nozzle is set at T<sub>3</sub>, and the temperatures are controlled such that the following relationship holds: T<sub>1</sub>, S T<sub>2</sub> S T<sub>3</sub>.
  - [0026] According to claim 2 of the invention, the substrate processing method described in Claim 1 is characterized in that the dry gas is obtained by bubbing the limit gas in the organic solvent, and the mixed gas is formed out of the intert gas including the organic solvent valve or containing the organic solvent was only solvent ways or containing the organic solvent mixet and the organic solvent ways.

concentration is lower than saturation point. The temperature of the mixed gas is controlled or maintained at constant temperature or gradually increased until the mixed gas is ejected from the jet nozzle. Therefore, the organic solvent is gradually veporized from the surface of the organic solvent mist as to decrease the diameter of the mist particles during movement, and the dry gas including the organic solvent mist of submisron size is easily obtained.

[0027] According to claim 3 of the invention, the substrate processing method described in claim 1 is characterized in that the dry gas which is a mixture of inert gas and the organic solvent rank or you believe the inert gas in an organic solvent in a vapor generating unit and is further diluted with dilution gas of the same kind of inert gas, wherein the temperature of the vapor generating unit is set at 11, the temperature of the mixed gas is set at 12, 'rom the vapor generating unit until the mixed gas is diluted with the dilution gas, the temperature of the dilution gas is set at 17, the temperature of the mixed gas containing the inert gas and the organic solvent is set at 12, to the jet nozzle after the mixed gas is diluted with the dilution gas, and the temperature of the dirute gas and the temperature of the dirute gas is directly and the temperatures are controlled such that the following relationship holds: 1, 5, 1, 5

[0028] According to the substrate processing method described in claim 3, the mixed gas formed in the vapor generating unit out of the inert gas and the organic solvent vapor containing the organic solvent mist and the organic solvent gas whose concentration is lower than saturation point, is further diluted with additional dilution gas of the same kind of inert gas used in the bubbling. The concentration of organic solvent vapor thereby decreases in the mixed gas while the possibility of condensation of the organic solvent mist diminishes, and the IPA mist can thus be conveyed to cover the spaces between the wafers. Therefore, since temperature of the mixed gas is controlled or maintained at constant temperature or gradually increased until the mixed gas is ejected from the jet nozzle, the efficiency and speed at which a part of the organic solvent is vaporized from the surface of the organic solvent mist to take the form of micro mist are increased, and a large amount of dry gas containing the organic solvent mist of submicron size can be obtained while the concentration thereof is low, thereby allowing a large quantity of the organic solvent mist of submicron size to be continuously injected to the substrate surface. Accordingly, even if multiple substrates of large diameter are loaded into the processing vessel, since the mist of submicron size can be rapidly applied to the spaces between substrates, the continuous supply of large amounts of organic solvent vapor of submicron size ensures the rapid substitution of the rinsing solution adhering to the substrate. As a result, processing time can be shortened while the level of efficiency of drying processing is improved, and acceleration of drying processing is achieved. Accordingly, processing time can be shortened while the level of efficiency of drying processing is improved, without increasing the amount of organic solvent to be consumed, and the generation of water marks on the substrate surface can be largely avoided or substantially eliminated. Further, adhesion of the particles is eliminated while accelerating the drying processing, so that re-adhesion

of particles can be prevented.

[0029] According to claim 4 of the invention, the substrate processing method described in any one of claims 1 to 3 is characterized in that the organic solvent is at least one kind's elected from a group including isopropyl alcohol, discretone alcohol; 1-methoxy-2-propanol, ethyl glycol, 1-propanol, 2-propanol, and tetrehydrofuran, and the inert gas is at least one kind selected from a rouse including introducen, aroon, and helium.

[0030] According to the substrate processing method described in claim 4, the selection width of the organic solvent and the inert gas is broadened, and the substrate processing apparatus can be applied to various kinds of substrate processing the any arbitrary combination.

[0031] The substrate processing apparatus according to claim 5 of the invention refers to a substrate processing apparatus comprising; a very generating unit which generates a mixture of an organic solvent twoprand an inert gas by bubbling the inert gas in an organic solvent; support means for supporting a plurality of substrates to be vertically arranged in parallel at equal pitches, a rinsing processing vessel which accommodates the plurality of substrates supported by the support means; a lid for covering the upper opening of the rinsing processing vessel; jet nozzles who are provided in the lid; and first piping which allows the vapor generating unit and the jet nozzles to communicate with each other, and the substrate processing apparatus is characterized in that the first piping and the jet nozzles are respectively equipped with heaters, and the heaters are controlled by means of dry gas containing organic solvent mists of submirrors size being emitted from the let nozzles.

[0032] According to the substrate processing apparatus described in claim 5, the dry gas including the organic solvent of submicron size can easily be formed by controlling the heaters which are provided at appropriate locations, and thus the substrate processing apparatus can easily carry out the substrate processing method described in claim 1.

[0033] According to claim 6 of the invention, the substrate processing apparatus described in claim 5 is characterized in that the temperature of the vapor generating unit is set at  $T_1$ , the temperature of the first pripring is set at  $T_2$ , and the temperature of the jet nozzle is set at  $T_3$ , the temperatures being controlled such that the following relationship holds:  $T_1 \le T_2 \le T_3 \le T_3$ .

[0034] According to claim 7 of the invention, the substrate processing apparatus described in claim 5 is characterized in that a second ciping is connected to the middle portion of the first piping for the purpose of suppying diffution gas of the same kind of inert gas, wherein the temperature of the vapor generating unit is set at T<sub>s</sub>, the temperature of the first piping is set at T<sub>s</sub> from the vapor generating unit to the point in which it is connected with the second piping, the

temperature of the second piping is set at  $T_4$ , the temperature in the first piping is set at  $T_2$ \* from the point in which it is connected to the second piping to the nozzle, and the temperature in the piperatures are controlled such that the following relationship holds:  $T_1 \leq T_0 \leq T_0 \leq T_0 \leq T_0$ .

[0035] The substrate processing apparatus described in either of claims 6 or 7 can easily carry out the substrate processing method described in either of claims 2 or 3.

[0036] According to claim 8 of the invention, the substrate processing apparatus described in claim 7 is characterized in that a static mixer is provided downstream from the point of connection between the first piping and the second piping and upstream in respect of the jet nozzle.

[0037] According to claim 8 of the invention, the substrate processing apparatus described in claim 7 is characterized in that the inert gas, the organic solvent mist, and the organic solvent mist, are said series with expectation in the state mixer is provided downstream from the point of connection between the first piping and the second piping and upstream in respect of the jet nozzle. Therefore, the formation efficiency of the micro mist is improved. [0038] According to claim 9 of the invention, the substrate processing apparatus described in any one of claims 5 to 8 is characterized in that the organic solvent is at least one kind selected from a group including isopropyl alcohol,

as is cherecterized in insit the origanic solvent is at least one knot section of more up including isopropy accions, discetone isotroni, I-methody-2-propanel, elity glycol, 1-propanel, 2-propanel, and tetrahydrofuran, and the inert gas is at least one kind selected from a group including nitrogen, argon, and helium.
(2021) According to the individual propagation of the intertion of t

[0039] According to the substrate processing apparatus described in claim 8, the selection width of the organic solvent and the inert gas is broadened, and the substrate processing apparatus can be applied to various kinds of substrate processing by any arbitrary combination.

#### BRIEF DESCRIPTION OF THE DRAWINGS

### [0040]

20

40

- Fig. 1 is a sectional view showing a substrate processing apparatus according to an embodiment of the invention;
  - Fig. 2 is a side view showing a processing vessel referred to in the embodiment of the invention;
  - Fig. 3 is a side view showing the processing vessel of Fig. 2 when viewed from the opposite side;
  - Fig. 4 is a plane phantom view of the processing vessel referred to in the embodiment of the invention when viewed from an upper portion of the lid thereof;
- 30 Fig. 5 is a side view of the lid illustrated in Fig. 4;
  - Fig. 6 is a view showing a timing chart of a series of processing steps;
  - Fig. 7 shows a cleaning and drying process, Fig. 7(a) is a sectional view illustrating the cleaning process, Fig. 7(b) is a sectional view illustrating the drying process 2, and
  - Fig. 7(d) is a sectional view illustrating the drying process 3;
  - Fig. 8 is a sectional view schematically showing the relationship between the IPA vapor and a substrate in the drying process;
    - Fig. 9 is a sectional view schematically showing the relationship between the IPA vapor and a substrate in the drying process as illustrated in Fig. 8, when dilution nitropen gas is used:
    - Fig. 10 is a sectional view showing the conventional substrate processing apparatus; and
  - Fig. 11 is a sectional view schematically showing the relationship between the IPA vapor and a substrate in the drying process of the substrate processing apparatus illustrated in Fig. 10.

### BEST MODE FOR CARRYING OUT THE INVENTION

- 49 [0041] A preferred embodiment of the invention will be described hereafter with reference to the drawings. However, the substrate processing method and substrate processing apparatus referred to in the embodiment are merely used as examples embodying the technical idea of the invention and are not intended to limit the applicability of the invention as the invention can be also applied to other modes without departing from the scope of claims.
  - [0042] Referring to Fig. 1, a substrate processing apparatus to its an apparatus for processing the semiconductor wafer W which represents an example of a substrate. The term 'processing' shall include the processes of etching the wafer W with chemicals, performing hydrofunds acid treatment to the surface of the wafer W, rinsing the wafer W, and applying drying processing to the wafer W after rinsing with the organic solvent, and the like. These processing steps are confinuously neutromed in series in one processing visual fig.
- [0043] As shown in Figs. 2 to 5, the processing vessel 15 is placed in a housing chamber 11 having sufficient capacity 5 with which to accommodate the processing vessel 15 and accessories thereof (not shown), which include an air conditioning device in the accommodation chamber, a supply source which supplies various processing solutions to the processing vessel, and a wafer conveying mechanism. The processing vessel 15 comprises an inner vessel 20, an outer vessel 25 and a lid 30. The inner vessel 20 an outer vessel 25 and the tuber surface thereof is exposed.

The outer vessel 25 surrounds the upper outer periphery of the inner vessel 20 and the opening of the inner vessel 20 is covered with the if 40. The inner vessel 20 and outer vessel 25, which are nade of material which is corresion-resistant to the organics ovent such as hydrollouric acid and IPA. e.g., polyvinylidene fluoride material, are eccommodated

[0044] The inner vessel 20 has sufficient depth to allow substrate processing by dipping a large number of waters W (e.g., about 50 waters W) having a diameter of a large as 000 mm in the processing solution white being held by a substrate holder 62 (see Fig. 3). A processing solution drain unit 21 and a processing solution supply unit 22 are provided on the bottom portion of the inner vessel 20. In the substrate holder 62, the plural waters W are vertically held in parallel by a cesseste guide (for example) at equal picties. The substrate holder 62 is outplet to a titing mechanism 60, which is provided with a lifting means 61, and the substrate holder 62 is taken out and put into the inner vessel 20 by vertically moving the substrate holder 62 with the #illing means 61. In Fig. 2, the drying process position is indicated as "Dry Position" while the rissing process position is indicated as "Rinse Position." An air cylinder mechanism may serve as the litting means 61.

[0045] As shown in Fig. 3, multiple wafers are taken out of the substrate holder 62 by means of a moving mechanism of 50. The moving mechanism So includes pursel holding pawls 50, and 50<sub>2</sub>, outget to a robot mechanism (not shown), and the wafers are held and moved to a pradetermined location by the holding pawls 50<sub>4</sub> and 50<sub>2</sub>. As shown in Fig. 2, the processing solution drain unit 21 is equipped with a noutlet 21<sub>1</sub> having a small diameter and an outlet 21<sub>2</sub> having a large diameter. The outlet 21<sub>2</sub> intentions as a draining mechanism which reptive discharges the processing solution the processing vessel. The outlet 21<sub>1</sub> having a small diameter drains the processing solution that has accumulated in the bottom portion of the inner vessel 20 and inside the piping. The outer vessel 25, which is provided with an outlet 25<sub>1</sub> at the lower portion thereof, functions as an overflow receptacle for the processing solution overflowing from the upper portion of the inner vessel 20.

[0046] As shown in Fig. 5, the lid 30 includes a box shaped container 31, the lower portion of which he exposed while the upper portion thereof is closed. Multiple wafers collected to form a group of wafers W can be housed in the box shaped container 31. The box-shaped container 31 is made of material which is corresion-resistant to the organic solvent suon as hydrofluoric acid and IPA. The lid 30 can be moved horizontally by moving means 55 (see Fig. 3). The opening of the inner vessel 20 can be closed and opened through the moving means 55, which causes the lid 30 to move horizontally on the upper portion of the inner vessel 20 as a shown in Fig. 2. In other words, the moving means 65 vertically lifts the lid 30 located on the inner vessel 20 for a predetermined distance, horizontally moves the lid 30, and then vertically sifts the lid 30 again to hold it in standby position. The lid 30 moves when the group of wafers W is conveyed

Into the inner vessel 20 or when they are taken out of the inner vessel 20 after processing.

[0047] As shown in Fig. 5, a ceiling surface 32 substantially having the shape of an arch is formed in the upper portion of the box-shaped container 31. In the ceiling surface 32, plural jet nozzles 33, to 33, which inject the linet gas are arrayed in four directions at substantially equal intervals. As shown in Fig. 4, in the plural jet nozzles 33, to 33, or carranged at substantially equal intervals, such six rows of jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such that the 42 jet nozzles 33, to 33, are arranged at substantially equal intervals, such manner that they are nearly equidistant from the outer circumference of the group of wafers. Since the wafer W is substantially disk-shaped, such distances can easily be made uniform by designing the ceiling surface 32 to have the shape of and. It is preferable that the shape of the ceiling surface be competitive with the shape of the wafer W to schieve substantial equality for such distances.

[0048] As shown in Fig. 5, a gas supply pipe 34<sub>p</sub> is connected to the jet nozzie 33, and the gas supply pipe 34<sub>p</sub> branches into branch pipes 44<sub>p</sub> and 34<sub>p</sub>. The same number of jet nozzies 33 to onsected to each of the branch pipes 84<sub>p</sub>, and 84<sub>p</sub>, and 84<sub>pp</sub>, and 84<sub>pp</sub>, and 84<sub>pp</sub>, and 84<sub>pp</sub> and 84<sub>pp</sub>

[0049] The jet nozzie 33 is besically conically shaped and provided with a hole on the tapered leading and thereof, from which the dry gas is ejected. Each jet nozzie 33 is equipped with a heater (not shown). Detailed description of the tonozzie with be omitted because it is publicly known. Further, the gas supply pile 94, and the branch pipes 34, and 34, and 34, and 34, are equipped with heaters (not shown) on the respective outer wall surfaces thereof. For example, a belt heater is used and the heaters are connected to a control processing unit (CPU) 12 which controls the heaters.

[0050] As shown in Figs. 2, 3, and 5, an intermediate connecting member 26 and a porous plate inserting mechanism

27 are arranged between the inner vessel 20 and outer vessel 25 and the lid 30. The intermediate connecting member 26 has a cylindrical body whose opening is the same as that of the lower opening of the lid 30, and is made of material which is corrosson-resistant to the organic solvent such as hydrofluoric acid and IPA. The intermediate connecting member 26 is arranged above the porous plate inserting mechanism 27 and a lower opening 25; is possitioned therein so as to about the upper surface of a frame body 27, in which a porous plate 8 is housed, and an upper opening 26; is fitted in the lower opening 31, of the box-shaped container 31. The intermediate connecting member 26 however may be omitted by directivit fitter the list 30 in the frame body 27.

[0051] The porous plate 28 includes a flat plate in which several small holes are made on the surface thereof. The flat plate is inserted between the inner vasses 20 and outer vessel 25 and the intermediate connecting member 26 during the process of drying the group of walkers W after the predetermined processes are finished. The porous plate 28, which is likewise made of material which is corrosion-resistant to the organic solvent such as hydrofluoric acid and IPA, is housed in the frame body 27, The porous plate 28 is coupled to a moving mechanism (not shown) and horborally moved in a sliding manner as shown in Fig. 2. The frame body 27, housing the porous plate 28 has a predetermined longitudinal width (in the vertical direction), and a gap 27<sub>2</sub> is formed between the frame body 27, and the porous plate 28 when the prorous plate 28 whe

[0052] The gap 27<sub>2</sub>, for example, is about 2 mm. In the drying process, part of the dry gas is outgassed into the sink 29 Therefore, since the gap 27<sub>2</sub> is formed between the inter vessel 20 and the lid 30 (pp 27<sub>2</sub> is surpressed by "x" in Fig. 7), a nid-foced state is formed between the inner vessel 20 and the lid 30 by the gap x, i.e., the half-closed state is formed between the drying processing unit and the risking processing unit and the sink 29. The porous plate 28 is inserted between the inner and outer vessels 20 and 25 and the intermediate connecting member 26 to partition the inner vessel 20 and the lid 30. In other words, the porous plate 28 functions as a shutter which separates the rinsing processing unit from the drying processing unit from the original processing unit and the state of the original processing unit and the state of the processing unit and the state

[0053] Hersafter, pcing connection between the processing vessel 15 and its accessories will be described with reference to Fig. 1. A processing solution introducing pipe 22, is connected to the processing solution supply visit as provided in the bottom portion of the inner vessel 20 and is also connected to a pure water supply source 38 through a flow control valve and a pump. The processing solution introducing pipe 22, together with the flow control and the pump serves as the processing solution supply system, which also includes the friends solution supply means. A chemical supply source 39 is also connected to the processing solution introducing pipe 22, through the flow control valve. The chemical supply source 39 includes chemical preparation means front shown for preparing the desired chemicals or predetermined concentration at a certain temperature. Depending on the purpose of processing for example, frieing, etching or oxidation, for example, the chemicals are selected from among hydrofloric acid, hydrochiofa aci

(0054) As shown in Fig. 2, the processing solution drain unit 21 provided in the bottom portion of the inner vessel 20 includes an outlet 21, of small diameter and an outlet 21, of large diameter. The outlets 21, and 212 are connected to the inner vessel drain pipes 23, and 23, respectively. The drain pipe 23, is connected to a drain processing facility 40 through an on-off valve, the pump, and the flow control valve. Similarly, the drain pipe 25 is connected to an exhaust processing facility 41 through the on-off valve, the pump, and the flow control valve. The sink 25 is also connected to an exhaust processing facility 41, A drain pipe 25, is connected to the lower portion of the outer vessel 25, and the drain pipe 25.

[0055] Further, a vacor supply mechanism 37, which is intended to accumulate the organic solvent such as isopropyl alcohol (IPA) solvent, is provided near the processing vessel 15. The organic solvent such as the IPA solvent easily mixes with the water remaining on and adhering to the surface of the water W, while having extremely little surface translor. The vapor supply mechanism 37 is provided with a vapor generation unil 37, which heats the organic solvent, vapor zes the organic solvent and generates mists by bubbling the inertigas in the organic solvent. The vapor generation unil 37, is dispect in hot water in a heating vessel 37, and the organic solvent is heated at predetermined temperature. The vapor generation unit 37, and the organic solvent supply source 36 are connected to each other with piping 36, and IPA is supplied to the vapor generation unit 37, Beadless IPA, the organic solvent must be selected from the group including organic compounds such as discetone alcohol, 1-methoxy-2-propanol, ethyl glycol, 1-propanol, 2-propanol, and terrahydrofuran.

[0056]. A second inert gas supply source 35 is connected to the veryor generation unit 37, through piping 35, and piping 35<sub>12</sub>. The nitrogen gas N<sub>2</sub> is supplied to the bottom portion of the vargor generation unit 37, to generate (bubbling) bubbles in IPA that has accumulated in the vapor generation unit 37, which forms the IPA vapor including the IPA gas and mist. Piping 37<sub>12</sub> derived from the vapor generation unit 37, is coupled to a gas supply pipe 34<sub>2</sub> through a static mixer M, and the mixed gas of the carrier gas N<sub>2</sub>, and the IPA vapor is supplied to the jet nozzle 33 from the vapor generation unit 37, Heaters (not shown) are respectively provided on the outer wall surfaces of the pipings 35<sub>12</sub>, 37<sub>12</sub>, and 34<sub>2</sub>, and the temperatures thereof are controlled by the CPU 12. Meanwhile, the static mixer M is provided in order to homogenize the mixed gas by accelerating the degree of mixture of the mixed gas containing the carrier gas N<sub>2</sub>.

the IPA vapor.

[0057] A first inert gas supply source 34 supplies inert gas such as nitrogen gas No through piping 34, to the piping 37;2. The temperature of the piping 34, is also controlled to a predetermined degree by the belt heater. The nitrogen gas No does not merely dilute the mixture of inert gas and the organic solvent vapor from the vapor generation unit 37. but is also useful for the purging and finish-drying steps in the processing vessel 15. Besides No. argon and helium are likewise appropriate for use as the inert gas.

[0058] Hereafter, the series of processes carried out by the substrate processing apparatus will be described with reference to Figs. 6 and 7. Referring to Figs. 1 and 6, first, the fid 30 of the processing vessel 15 is opened, and multiple wafers W are accommodated in the inner vessel 20, which is supplied at this point with the desired chemicals, e.g., hydrofluoric acid (HF) from the chemical supply source 39 through the processing solution introducing pipe 22, and the processing solution supply unit 22, and the desired chemicals are then stored in the inner vessel 20, Accordingly, substrate processing (for example, etching, hydrofluoric acid treatment, and rinsing) is performed to the wafers W by dipping them in the processing solution.

[0059] Upon the completion of such chemical processing, as shown in Fig. 7(a), the pure water DIW is supplied to the inner vessel 20 from the pure water supply source 38 through the processing solution introducing pipe 22, and the processing solution supply unit 22, while overflowing from the upper portion of the inner vessel 20. The overflowing pure water DIW flows into the outer vesse: 25, and is drained from the drain pipe 25, through the drain pipe. The pure water is supplied for a relatively long period of time to wash out the chemicals HF which have remained in the inner vessel 20. [0060] After the rinsing process, in one particular drying process 1 illustrated in Fig. 7(b), the continuous supply of pure water DfW is terminated, and the group of wafers W is slowly raised from the inner vessel 20 (slow up speed) while being supplied with a small amount of pure water (DIW water-saving). The small amount of IPA vapor can also be simultaneously supplied into the processing vessel 15 while the group of waters W' is being raised.

[0061] Then, in another drying process 2 illustrated in Fig. 7(c), a drain mechanism valve of the cutlet 212 lying at the bottom portion of the processing vessel 15 is operated to drain the processing solution rapidly, and the porous plate 28 is inserted between the inner vessel 20 and outer vessel 25 and the intermediate connecting member 26 by causing the porous piate 28 to horizontally move into the frame body 27, Further, the hot dry gas containing the mixture of nitrogen gas No and the IPA vapor is supplied to the inner vessel 20. These operations are performed as shown in the chart.

[0062] The dry gas is heated in the piping 34, and the nozzle 33. In the drying process 2, the organic solvent vapor in the processing vessel 15 comes into contact with the surface of each wafer W, and the IPA mist is condensed onto the surface of the wafer W to form a film of IPA. When the organic solvent film is formed on the surface of the wafer W, IPA is substituted for the pure water adhering to the water W. Therefore, the degree of surface tension of the pure water decreases to allow it to flow down from or evacuate the surface of the wafer W, and IPA adhering to the substrate surface is evaporated.

[0063] In another drying process 3 illustrated in Fig. 7(d), the nitrogen gas N<sub>a</sub> is supplied in order to dry the substituted IPA, and the group of wafers W' is taken out of the processing vessel 15 when the drying process 3 ends.

[0064] In the above processes, pressure in the fid 30 (the drying processing unit) is set higher than that of the sink and the evacuating station for the sink, and pressure in the lid 30 is also set higher than that of the inner vessel 20 (rinsing processing unit) and pressure of the evacuating station for the inner vessel 20. Therefore, the nature of the flow of the dry gas becomes laminar flow in the drying processing unit, and the dry gas is smoothly evacuated from the exhaust pipe to the outside of the vessel. During this process, the dry gas is evenly supplied to each of the waters, water marks do not form on the substrate surface, and particles that have settled thereon can be removed, and particles can be prevented from adhering to the substrate surface. Further, the re-adhesion of particles can be avoided because the dry das does not circulate at all in the processing yessel.

[0065] At this point, the CPU 12 controls the heaters such that the relationship among the temperatures T<sub>1</sub>, T<sub>2</sub>', T<sub>4</sub>, T2", and T3 at certain positions satisfies the following relationship:

$$T_1 \le T_2' \le T_4 \le T_2'' \le T_3$$
 (1)

where

the temperature in the vapor generating vessel 37, is T<sub>1</sub>.

the temperature in the piping 3742 is T2',

the temperature in the piping 34, is Tax

the temperature in the piping 342 is T2\*, and

the temperature in the jet nozzie is T<sub>3</sub>.

[0066] Thus, the temperatures T<sub>1</sub>, T<sub>2</sub>', T<sub>4</sub>, T<sub>2</sub>', and T<sub>3</sub> at respective positions are set so as to satisfy the above expression (1), and thereby allow the size of the IPA mist emitted from the let nozzle 33 to have an extremely small

particle diameter, or specifically, extremely fine mist which cannot be observed in relation to the Tyndall phenomenon, i.e., mist having submicron size is formed. In other words, the mist can be observed by the Tyndall phenomenon if it consists of several particles having micron size but cannot be seen in relation to the Tyndall phenomenon when their comprises particles of submicron size, thereby requiring the use of a particular measuring device. In the embodiment of the present invention, since the size of the IPA mist cannot be confirmed by the Tyndall phenomenon, the mist formed constitutes particles of submicron size. However, it has been noted that the mist of submicron size is not completely formed in the gas, but exists in the liquid state. The temperature in the jet nozzle or T<sub>g</sub> is controlled so as not to exceed the obling point (32.4 °C) of IPA to prevent the IPA mist from being completely apparature.

[0067] The number of particles of the organic solvent mist can be increased without increasing the amount of organic solvent to be consumed by ensuring that the mist constituting the organic solvent vapor is of submirrors size. Further, although the surface area of the individual mist is reduced, the whose surface area which is the summation of the surface areas of the particular mists proportionately increases in relation to the increase in the number of mists. As a result, because the organic solvent vapor can be injected to the substrate while a specific surface area thereof is increased, the level of substitution afficiency of the organic solvent for the rising solution is improved.

[0068] It is preferable that each temperature referred to in expression (1) is kept constant or gradually increased. That is, the mixed gas formed out of the inert gas including the IPA vapor containing the IPA mix and the IPA gas whose concentration is lower than the point of saturations is obtained by bubbling the inert gas in the IPA in the vapor generation until 37. The temperature of the mixed gas is kept constant or gradually increased until it is emitted from the jet nozzle, so that the condensation of organic solvent vapor in the piping and the jet nozzle is eliminated and as a result, the diameter of the organic solvent mixt does not grow in the mixed gas. Further, IPA is gradually vaporized from the surface of the IPA mixt to decrease the particle diameter of the IPA mixt while the IPA moves from the piping and the nozzle, thereby allowing dry cas containing IPA mixer of submirrom size to be easily obtained.

[0069] Further, the mixed gas formed in the vapor generation unit 37, out of the inert gas including the IPA vapor containing IPA mists and the IPA gas whose concentration is lower than the point of saturation is disluded with inert gas separately supplied from the piping 34,. Therefore, since the concentration of IPA gas diminishes in the mixed gas, dry gas containing IPA mists of abunitorion size can be supplied to the processing vessel 15 along with a large amount of inert gas while IPA vapor.atdon from the IPA mist is excelerated, in this case, in mixing the newly supplied introgon S. N, with IPA vapor and the IIIA, it is preferable that a mixer M is provided in the middle of the piping 34, to stir the nitrogen sas. N, and the IPA vapor. A static mixer is suitable as mixer M.

(0070) Thus, when numerous IPA mists of submicron sizes are contained in the dry gas, such mists can rapidly invade the spaces between substrates en if many substrates having a large diameter are loaded in the processing vessel 15, and the rapid supply ocontinuously large amounts of organic solvent vapor of submicron size to replace the rinsing solution atthering to the substrate is assured. As a result, substrate processing time can be shortened while the rate of drying processing efficiency is simultaneously improved, and the remarkable acceleration of the drying processing can be achieved. At the same time, the amount of organic solvent need not be particularly increased, while the generation of watermarks can be extremely reduced or substantially eliminated in the substrate surface. Further, particle adhesion is eliminated and the drying processing is a societized so that re-achesion of particles can be also prevented.

[0071] In the above mode, the linert gas for dillution is supplied from the piping 34. The inert gas for dillution is not always required in order to obtain IPA mists of submicron size. In this case, temperature or control shriply may be performed on as to satisfy the expression (1), while the temperatures of the piping pieces 373, a 43, 434, and 344<sub>2</sub>, which connect the vapor generation unit 37, and the nozzle 33 differ from one another. Alternatively, the pieces of piping 37<sub>12</sub>, 34<sub>2</sub>, 34<sub>2</sub>, and 34<sub>22</sub> are kept at the same temperature T<sub>2</sub>, and temperature may be controlled such that the following relationship holds:

# $T_1 \leq T_2 \leq T_3.$

[0072] The temperature T<sub>3</sub> in the piping and the jet nozzle is set equal to or higher than the temperature T<sub>3</sub> of the organic solvent and the temperature T<sub>3</sub> of the interfigus supplied to the velop repensating unit. In this meanure, condensation of the organic solvent vapor in the piping and the jet nozzle does not occur, so that the dismeter of the organic solvent mist in the dry gas does not grow. In addition, part of the organic solvent is further vaporized from the surface of the organic solvent mist before the organic solvent is before the organic solvent is the control to the organic solvent is entitled from the nozzle, which allows the size of the organic solvent mist in the dry gas to become submicron. Further, since the organic solvent vapor is not condensed in the piping and the jet nozzle, there is no learn that IPA droplets will fall from the nozzle.

[0073] In the embodiment of the invention, it is preferable that the components of the substrate processing apparatus are configured as follows, in order to prevent the increase in diameter of the organic solvent mist and condensation thereof due to the collision of the contains solvent mist with each other and the collision of the contains solvent mists with each other and the collision of the contains solvent mists with

the walls of the apparatus. That is, the top portion of the bubbling tank of the vapor generating unit should have a conical or circular are shape. In addition, the number of step differences in the dry gas supply piping has been decreased, and apertures are formed such that the size thereof is not remarkably bigger than that of prior art. Further, the bubbling nozzle in the vapor generating unit should have a small diameter and quick injection speed.

[0074] Finally, the state in which the IPA mist is made to substitute for the rinsing solution adhering to the substrate by using dry gas containing IPA mists of submicron size will be described. Figs. 8 and 9 are sectional views schematically showing the relationship between the IPA vapor and the substrate during the drying processing. Fig. 8 illustrates the case where the dry gas is not diluted with inert gas white containing IPA mists of submicron size, while Fig. 9 shows the case in which the dry gas is diluted with linert gas white containing IPA mists of submicron size, while Figs. 8(a) and 8(b) and Figs. 9(a) and 9(b) correspond to the process of "drying 2" shown in Fig. 6, while Figs. 8(c) and Fig. 9(c) correspond to the process of "drying 2" shown in Fig. 6.

[0075] As shown in Fig. 8(a), when the dry gas containing the mixture of iPA mists (liquid) of submicron size and the ntrogen gas N<sub>2</sub> (gas) is delivered to the processing vessel 15 and supplied in the spaces between the wafers W. IPA is substituted for DIW adhering to the wafer W lyth supply of dry gas as shown in [16, 8(b). Since the IPA mist comprises

In particles and many IPA mists are supplied, various IPA particles adhere to one DIM, and the IPA mist therefore efficiently substituted for DIW. Then, as shown in Fig. 8(c), IPA is evaporated only by delivering N<sub>2</sub> gas into the processor in great processor in Fig. 8(a) and 8(b), the IPA mist does not sufficiently perform the carrier effect due to the small amount of nitrogen gas N<sub>2</sub>, supplied and good drying effect is obtained only to some degree by using IPA mists of submicron size. Still, however, on occasion water marks are generated by residual DIW in the process of drying waters with a large diameter.

(1076) In the case where the dry gas conteining IPA mists of submicron size is further diluted with inert gas, the IPA vapor concentration in the dry gas is reduced, such that the IPA mist is sufficiently distributed throughout the substrate. Therefore, as shown in Fig. 9, the IPA mists of submicron size can be continuously and evenly supplied even to deeper portions of the wafers W due to the large amounts of carrier gas. Since the dry gas containing large amounts of carrier gas enables the IPA mists of submicron size to adhere repully to DIW, efficient substitution of DIW by the mists takes place. Accordingly, the invention may also be used for drying waters having a large diameter, since the level of drying processing efficiency has been increased, while reducing processing time because processing speed is accelerated. Enrich, as shown in Fig. 9(c), DIW can substantially be eliminated from the surface of the water W, such that the generation of water marks can be completely eliminated. A detailed description of Fig. 9 has been omitted since it differs from the case shown in Fig. 8 only in that new nitrogen gas is mixed.

### Claims

- 39 1. A substrate processing method in which the surface of a substrate is dried by injecting it with dry gas consisting of a mixture of an organic solvent vapor and an inert gas, characterized in that the organic solvent vapor contains mists of submirron size.
- A substrate processing method according to claim 1, characterized in that the dry gas consists of a mixture of inert gas and the organic solvent vapor, being formed by bubbling the inert gas in the organic solvent in a vapor generating unit;
  - wherein the temperature of the vapor generating unit is set at  $T_t$ , the temperature of the mixed gas containing the inert gas and the organic solvent is set at  $T_2$  from the vapor generating unit to a jet nozzle, and the temperature of the dry cas emitted from the jet nozzle is set at  $T_2$ , and
- 45 the temperatures are controlled such that the following relationship holds:

# $T_1 \leq T_2 \leq T_3.$

- A substrate processing method according to claim 1, characterized in that the dry gas consisting of a mixture of finer gas and the organic solvent vapor is formed by bubbling the linert gas in the organic solvent in a vapor generating unit, and is further diluted with dilution gas of the same kind of inert gas.
- wherein the temperature of the vapor generating unit is set at T<sub>x</sub>, the temperature of the mixed gas is set at T<sub>x</sub> from the vapor generating unit until the mixed gas is diluted with the dilution gas, the temperature of the dilution gas is set at T<sub>x</sub>, the temperature of the mixed gas containing the linet gas and the organic solvent is set at T<sub>x</sub> to the jet nozzle after the mixed gas is diluted with the dilution gas, and the temperature of the dry gas emitted from the jet nozzle is set at T<sub>x</sub> and

the temperatures are controlled such that the following relationship holds:

# $T_1 < T_2' < T_4 < T_2'' < T_3$ .

- 4. A substrate processing method according to any one of claims 1 to 3, characterized in that the organic solvent is at least one kind selected from a group including isopropy! alcohol, diacetone alcohol, 1-methoxy-2-propanol, ethyl glycol, 1-propanol, 2-propanol, and tatrahydrofuran, and the inert gas is at least one kind selected from a group including nitrogan, argon, and helium.
- 5. A substrate processing apparatus including:
  - a vapor generating unit which generates a mixture of organic solvent vapor and an inert gas by bubbling the inert cas in an organic solvent:
    - support means for supporting a plurality of substrates to be vertically arranged in parallel at equal pitches;
  - a rinsing processing vessel which accommodates the piurality of substrates supported by the support means; a lid for covering the upper opening of the rinsing processing vessel;
  - let nozzles which are provided in the lid; and
- first ploing which allows the vapor generating unit and the jet nozzles to communicate with each other, the substrate processing apparatus **characterized in that** the first piping and the jet nozzles are respectively
  - equipped with heaters, and the heaters are controlled by means of dry gas containing organic solvent mists of submicron size being emitted from the jet nozzles.
  - 6. A substrate processing apparatus according to daim 5, characterized in that the temperature of the vapor generating unit is set at T<sub>1</sub>, the temperature of the first piping is set at T<sub>2</sub>, and the temperature of the jet nozzle is set at T<sub>3</sub>, and the temperatures are controlled such that the following relationship holds:

$$T_1 < T_2 < T_3$$
.

- A substrate processing apparatus according to claim 5, characterized in that the second piping is further included by being connected to the middle of the first piping for the purpose of supplying dilution gas of the same kind of lined gas.
  - wherein the temperature of the vapor generating unit is set at  $T_1$ , the temperature of the first piping is set at  $T_2$  from the vapor generating unit to the point in which it is connected with the second piping, the temperature of the first piping is set at  $T_2$ , from the point in which it is connected with the second piping is the said nozzle, and the temperature of the first piping is set at  $T_2$ , from the point in which it is connected with the second piping to the said nozzle, and the temperature of the jet nozzle is set at  $T_3$ , and the temperatures are controlled such that the following relationship holds:

# $T_1 \le T_2' \le T_4 \le T_2'' \le T_3$ .

- A substrate processing apparatus according to claim 7, characterized in that a static mixer is provided downstream from the point of connection between the first piping and the second piping and upstream in respect of the jet nozzle.
- 9. A substrate processing apparatus according to any one of claims 5 to 8, characterized in that the organic solvent is at least one kind selected from a group including isopropyl alcohol, disacetone alcohol, 1-methoxy-2-propanol, ethyl glycot, 1-propanol, 2- propenol, and tetrahydrofuran, and the inert gas is at least one kind selected from a group including nitrogen, argon and helium.

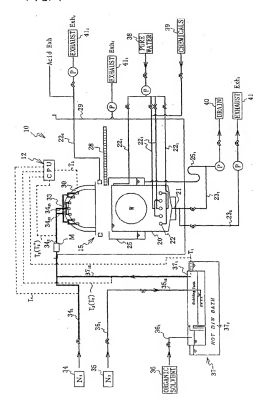
125

15

30

40

FIG. 1



F I G. 2

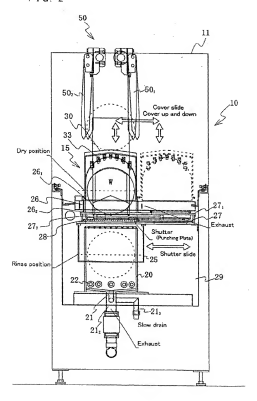
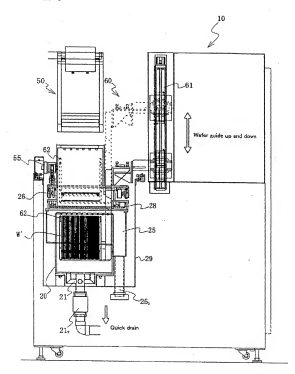


FIG. 3



F I G. 4

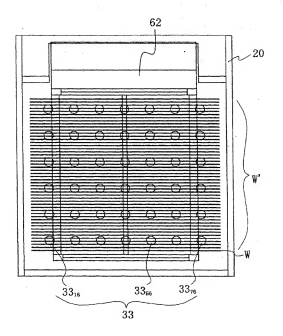


FIG. 5

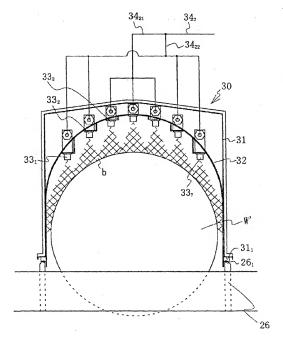
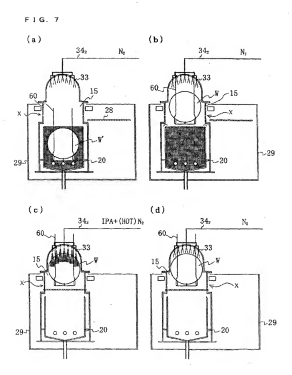
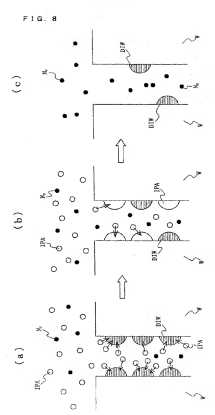


FIG. 6

	Rinsing	Dryingl	Drying2	Drying3
HF Dip			:	
MIG				
DIW .				
Slow up Speed				
Purge N:				
- Quick Drain			$\overline{\Box}$	
Porous plate D			n T	-
IPA supply			N2 IPA	ĺ
Dry No			,,,,	





F1G. 9

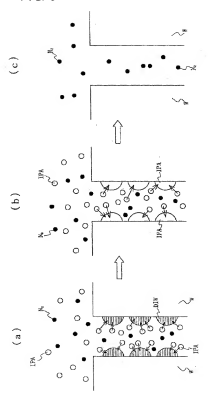
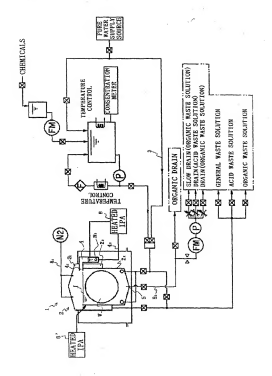
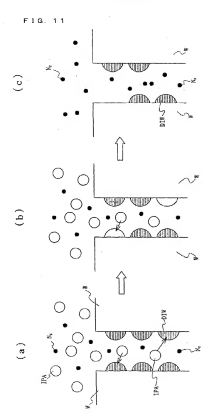


FIG. 10





# INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/15431

A. CLAS	SEFICATION OF SUBJECT MATTER C1 H01L21/304		
According	to International Patent Classification (IPC) or to both	national classification and IPC	
B. FIELD	S SEARCHED		
Minimum d Int.	ocumentation searched (classification system tollower C1	l by classification symbols)	-
Vits Koka	tiou serched other Nem ministerium documentarion to it iyo Shinan Koho 1925-1996 i Jitsuyo Shinan Koho 1971-2004	Toroku Jitsuyo Shinan Kok Jitsuyo Shinan Toroku Kok	ю 1994-2004 ю 1996-2004
	late frace consulted during the international search (na	me of data base and, where practicable, sa	nch terms used)
	MENTS CONSIDERED TO BE RELEVANT		
Category*	Clution of document, with indication, where a		Relevant to claim No.
Ă	JP 2000-55543 A (Tokyo Elec 25 February, 2000 (25.62.00) Full text; Figs. 1 to 10 (Family: none)		1-9
λ	JP 2002-134461 A (Sony Corp 10 May, 2002 (10.05.02), Full text; Figs. 1 to 11 (Family: none)	.), ·	1-4,6-9
¥	JF 10-210586 A (Komatou Itd. 12 May, 1998 (12.05.98), Full text; Figs. 1 to 5 (Family: none)		14,69
Purch	er documents are listed in the continuation of Box C.	See patent family agues.	
"A" decume constitle carbor o date "12" decurse cited to openial docume means docume than the	categories of doted documentaries and officially designed asset of the art which is not only to the opportunities abstract on the art which is not only to the opportunities abstract on or where the international filling with which are any through complete and the opportunities are also as the opportunities of a contract of the opportunities of a contract of the opportunities of a contract of the opportunities of the opportuni	"I" later documents profitible of their time in an periody does not exist a condition with it periody does not exist a condition with it "I" when the condition of the condition of considered more due of control to consider considered more due of control to consider "I" see when the document is taken above considered with case or more other con- considered to length as an exempt of considered to the control of the con- considered of the international season 2.4 February, 2004.	a application had cled to advise the invention accused be not been a considered desired inventions accused be not been a considered accusation accused be order the decomment in decomment, such askilled in the art husby
Name and mailing soldness of the ISA/ Japanese Patent Office		Authorized officer Telestone: No.	•
Facsimile No			

Form PCT/ISA/210 (second sheet) (July 1998)

# INTERNATIONAL SEARCH REPORT

International application No. PCT/JP03/15431

ategory*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim N
Y	JF 11-191549 A (Dainippon Screen Mfg. Co., Ltd.), 13 July, 1999 (13.07.99), Pull text: Figs. 1 to 4 (Family: none)	3,7
¥	JP 2002-359221 A (M-PSI Kabushiki Kaisha), 13 December, 2002 (13.12.02), Pull toxt, Figs. 1 to 15 (Pamily: nome)	8
***************************************		
	*	
***************************************		
***************************************		
-		
***************************************		
	-	

Figur PCT/ISA/210 (continuation of second sheet) (July 1998)